SPECIFICATION PATENT

NO DRAWINGS

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COMPLETE SPECIFICATION

Production of Heptafluoropropane

We, Allied Chemical Cobroration, a Corporation organised and existing under the laws of the State of New York, United States of America, of 61, Broadway, New York 6, New York, United States of America, do hereby declare the invention for mixing the present that he counted which we pray that a patent may be stunted to us and rise method by which it is to be performed, to be particularly described in and by the following statement:

This invention relates to the production of 111 2 2 2 2 harmfluorescences.

This invention relates to the production of 1,1,1,2,3,3,3-hepsalluotopropane, CF, CFH-CF, bp. -17 so -18.5 C., (hereinafter referred to simply as imptalluotopropane) by direct hydrofluormation of hexafturopropene, CF, CF = CF₂, bp. -29° C. Hepsalluotopropane is useful as a propellant and as a creater distance. gascous dielectric.

It has been proposed to make heptafluoro-propane by a procedure involving liquid-phase reaction between hexaliuoroprofene and aqueous hydrodiuoriu acid in a pressure vessel substantial superarmospheric pressures.

The disadvantages of batch, liquid-phase,
autoclave operations are apparent.

It has now been found that activated carbon

by itself possesses the pawer of promoting reaction of aphydrous hydrogen fluoride and hexafluoropropene in an eaxily controllable, all gas-phase, catalytic process in which heptatheoropropane is made in high yields. It is also of major importance, from the It is also of major importance, from the standpoint of simple operation, that activated carbon promotes the reaction at substantially amongharise pressure, and requires only moderately elevated reaction temperatures.

Generally, the practice of the invention includes effecting reaction between gaseous hexallucropropene and gaseous substantially aphydrous indrocer fittendic at moderately

anhydrous hydrogen finoshie at moderately annyorous ayurogen monate at moustainy statistics and in the statistics of activated carbon at satisfy, and presence of activated carbon at satisfy, and recovering heptafluoropropene from the re-sulting reaction products. More particu-

larly, gaseous hexaflueropropene and gaseous anhydrous hydrogen fluoride may be continuously metered, mixed and fed at substantially atmospheric pressure into a tubular reactor which is packed preferably full with activated carbon catalyst and which may be used of inert restricted made as which be made of inert material such as mickel, "Mcnui", or "Income" or of steel lined with graphic or "Alundum", (the words "Monei", "Incomel" and "Ahundum" being Registered Trade Marks), and enveloped in a suitable multar electric furnace provided with euromatic heating means to provide the reaction temperature. Product recovery may be Product recovery may be matte heating meants recovery may be emperature. Product recovery may be effected by means generally conventional in this art. For example, vapours leaving the reaction zone may be passed through a water scrubber to remove hydrogen fluoride, and those leaving the saudber may be dried and then totally condensed in the receiver middle condings such as by use of a and then totally condensed in the receiver by suitable cooling, such as by use of a lary lee-accepte mixture. The resulting condensers that may be fractionally distribed to freditate recovery of the sought-for product, and separation of possibly small amounts of unreacted starting material which may be recycled if desired.

The activated cartain caralysts which are

The activated carbon catalysts which are used in the practice of the invention may be used in the practice of the invention may be gratular materials which are readily available from several commercial sources, usitable materials being various grades of around \$2.14 mesh activated carbon such as Columbia 6G, Columbia SW, and Darco. (The word "Darco" is a Registered Trade Mark.) The grain size is not highly critical. Ordinarily, the reaction is carried out in elongared subular regions, and it is then desirable to employ reactors, and it is then desirable to employ activated earlier or grantles of average most size between 1/20th and 1/4th of the reactor diameter; preferably the reactor is sub-stantially completely filled with granules of average mesh size of about 1/8th or 1/10th of its diameter.

In order to secure the unusually high yields 90

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obtainable by the practice of the invention, it has been found that the reaction zone temperature should be held in the range of 250—450° C. Experience indicates that the best yields appear to be dependent upon a cerrelation of temperature and mol ratio of hydrogen fluoride to herafinerepropene sturing material. The reaction involved may be represented by

CF-CF=CF₀+HF -> CF-CFHCF.

Generally, the quantity of hydrogen fluoride used relative to herafinerepropene starting material may be any amount sufficient to

material may be any amount sufficient to react with a substantial quantity of the hexaflucropropens, and if hexaflucropropens conversion and utilization of hydrugen conversion and utilization of hydrogen fluoride per pass are not of major importance, hydrogen fluoride in amount less than theoretical may be employed. However, for 20 better overall operation, the mol ratio of hydrogen fluoride to hexaficoropropens preferably should be in the range of 1:1 to 3:1. When high yields are desired, it is preferred to employ the hydrogen fluoride in amount in cross of the theoretical requirement, and for best yields (of or approaching 100% on the basis of the erganic starting material feet) the basis of the organic starting material fed) a mol ratio of bydrogen fittoride to hexafluoropropene in the range of 1.5:1 to 3:1 is preferred. As demonstrated in the appended examples, when utilizing hydrogen-fluoride and organic starting material in the nuoride and organic starting material in the preferred mid ratio range, and employing reaction zone temperatures in the range of hepuffuoropropane may be obtained.

A major advantage of the invention is that the reaction may be carried our efficiently at the reaction may be carried our efficiently at the reaction of the control of the control of the carried our efficiently at the carried our efficiently at

the reaction may be carried our efficiently at substantially atmospheric pressure as distinguished from the superatmospheric pressures characteristic of the prior str. It will be understood that in the practice of graphase caulytic processes of the general type described herein, i.e. processes in which a gas stream is caused to flow successively through reaction and product recovery systems, the pressure is considered for practical purposes as being substantially atmoscal purposes as being substantially atmospheric even though it is sufficiently high on the input side to effect commercially satisfactions of the input side to effect commercially satisfactions. the input side to effect commercially statisticity gas flow through the apparatus train.

Thus, strictly speaking, depending upon factors such as apparatus design, mesh size of catalyst, unpacked gas space in the reactor, desired contact time, etc., the actual pressures in systems of the kind under consideration may vacy from 2 up to say 10—15 pounds gauge, and accordingly operations of this type are regarded as being effected under

type are regarded as being effected under substantially atmospheric pressure. Contact time may be varied considerably without noticeable falling off. In process efficiency. Concrally, an increase in the contact time and reactor temperature results in higher utilization of the hydrogen fluoride

and conversion of hexafluoropropene, and a decrease in contact time and reactor temperature results in lower unbigation of hydrogen fluoride and lower conversion. Contact times may lie in the range of 0.5—60 serronds, times may be in the range of U.5—60 serroids, but will usually and preferably be in the range of 1—25 seconds. To a substantial extent, constact time, reactor temperature and ratio of reactions are inter-related, and depending upon the apparatus and the particular operation at hand, optimum conditions as to contact wine may be determined by test runs.

In Examples I and II which follow, the reactor consisted of a 1 both internal diameter nickel tube 42 inches long heated externally nickel tube 42 inches long heated externally by an electric furnace enveloping about 30 inches of its length. The reactor was provided at the infect end with suitable means for metered introduction of greeous hexaftuoriopropure and anhydrous hydrogen fluoride, while the reactor outlet was connected to the bales and of a preducts recovery nected to the inlet and of a products recovery train. The camiyst employed consisted of activated carbon (commercially available Columbia 6G grade) of 8—14 mesh, the size being such as to provide granules averaging about 1/10th of the reactor diameter. The reactor was complemely filled with catalyst, the total votume amounting to about 0.54 little. The pressure in the reactor system was about 3 th. psig. i.e. sufficient to maye the gas attemn through sufficient to move the gas stream through the reactor system at the contact times indicated and through the remainder of the 100 apparatus train. Percentages are by weight.

HILUPIR T During a period of about 125 minutes, about 77 grems (3.85 mols) of suhydrous hydrogen filtoride and about 308 g. (2.05 to bydrogen filtoride and about 308 mals) of hexaffuorepropens were premixed and metered into the reactor system.

Throughout the run the temperature within system. the reactor was maintained in the range 392 to 402° C. Overall contact time was approximately 9 seconds. The vapours Icaving the resetor were water-scrubbed to remove some hydrogen fluoride, dried by passage through a calcium chioride drying tower, and unidensed and collected in a receiver confed with Dry Ice-actions. A total of 36.1 g. (1.81 mols) hydrogen fluoride was scrubbed out of the reactor exit gas. A total of 342 g. of material was condensed and recovered in the receiver. On distillation of the condensate, there were recovered 316 or the condensate, there were recovered 316 g of marental boiling in the range -16 to -17.5° C, and 25 g, of still residue. Analysis, including infra-red absorption spectrum analysis and gus chromatography, established both distillate and residue, a total of 341 c 17.0 m. 21 c. 18.0 m. of 341 g. (2.0 mola), to be hepathuoropropere. Hence, the yield on the beam of hemiticoro-propene fed was substantially 100%.

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Onclog a period of about 120 minutes 88 g. (4.40 mols) of anhydrory hydrogen thurside and about 278 g. (1.85 mols) of bernstoropropose were premised and meterod into the reason. Throughout the ron the temperature in the present was assumined in EXAMPLE II berafinoropropene were premixed and meterod into the reactor. Throughout the run the temperature in the reactor was maintained in the range 300 to 306° C. Overall contact the range as in Example I. A total of about 43 g. (2.15 mols) of hydrogen fluoride was scrubbed out of the reactor exit 528, and a total of out of the reactor exit 528, and a total of out of the reactor exit 528, and a total of out of the reactor exit 528, and a total of out of the reactor exit 528, and a total of out of the reactor exit 528, and a total of out of the reactor exit 528, and a total of the strap. On distillation there were recovered as condensate about 291 g. of material boiling submanishly in the range of -16 to -17.5° C. and 25 g. of still recidude. Analysis, including infrared about particular analysis and gas chromatography, showed both distillate and residue to be hepisiluopopropene, in total thread on hexafluoropropene fed was substantially 100%.

WHAT WE CLAIM IS:

1. Process for the production of 1,1,1,2,-

1. Process for the production of 1,1,1,2,-3,3,3-heptalluoropropone, which comprises bringing a mixture of herafinoropropeus and substantially arhydrous hydrogen funcide in the gas phase into contact with an active

curbon catalyst at a temperature of 250°-450°

2. Process according to Claim 1, wherein the reaction is effected under substantially

amospheric pressilit.

3. Process according to Claim 1 or 2, wherein the mol ratio of the hydrogen fluoride to the hemitioropropene is between 1:1

4. Process according to Claim 3, wherein the mol ratio is between 1.5:1 and 3:1 and 3:1 and special reaction temperature is 375 425° C.

the reaction temperature is 375°—425° C.

5. Precess according to any one of the preceding claims, wherein the mixture of hydrogen flooride and hearimoroproperse is fed continuously into and through a unfular reaction zone containing the active carbon and heated to the reaction temperature, and the versours feating the zone are continuously treated to isolate heptafluoropropane therefrom.

6. Process for the production of 1,1,1,2,-3,3,3-hepminoropropare actualing to Christ 1, substantially as hereinbefore described.
7. 1,1,1,2,3,3,3-hepminoropropane obtained by a process claimed in any one of the process claims.

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